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jc714 U.S. PTO



02/16/00

PROVISIONAL APPLICATION COVER SHEET

This is a request for filing a PROVISIONAL APPLICATION under 37 CFR 1.53(b)(2)

AIPRAW

Docket Number		11004-1		Type a Plus Sign (+) inside this Box -->		+
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TITLE OF THE INVENTION (280 CHARACTERS MAX.)						
Oil Content And Composition enhancement by Targeted/Non-Targeted Expression Of Plant Plastidial And Bacterial Gene For Glycerol-3-Phosphate Acyltransferase						
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STATE	Ontario	ZIP CODE	K1A 0R6	COUNTRY	Canada	
ENCLOSED APPLICATION PARTS (Check all that Apply)						
X	Specification	Number of pages	8	Small Entity Statement		
X	Drawing(s)	Number of Sheets	6	Other (specify)		
METHOD OF PAYMENT (Check One)						
	A check or money order is enclosed to cover the Provisional filing fees			Provisional Filing Fee Amount (\$)	\$ 150.00	
X	The Commissioner is hereby authorized to charge filing fees and credit Deposit Account Number 14-0429					

jc526 U.S. PTO
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02/16/00The invention was made by an agency of the United States Government or under contract with an agency of the United States Government. ☒ No

Date:

17 Feb 2000

Respectfully submitted,

J. Wayne Anderson
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Regn No. 28,158Enclosures
:ech

Additional inventors are being named a separately numbered sheets attached hereto.

PROVISIONAL APPLICATION FILING ONLY

**Oil Content and Composition Enhancement by Targeted/Non-Targeted
Expression of Plant Plastidial and Bacterial Gene For Glycerol-3-
Phosphate Acyltransferase**

BACKGROUND OF THE INVENTION

There is a general industrial demand for higher oil content in living organisms in order to be more cost effective during production and processing. It is estimated that a 1% increase in seed oil would add \$20 M to the value of the Canadian canola crop at the farm gate.

One way of accomplishing it is by modifying metabolic pathways of living organisms by enhancing the expression of genes involved in oil synthesis. For example, overexpression of a yeast *SLC1-1* gene has been found to increase oil content in plants (Zou et al., 1997).

Only a limited number of genes have been reported to enhance oil content (for example, Zou et al., 1997, Marillia et al., 1998). There is a need to explore other approaches using other genes which might be more effective alone or in combination with previously cloned genes.

SUMMARY OF THE INVENTION

We propose to increase seed oil content and modify seed oil composition by overexpressing the genes for the enzyme glycerol-3-phosphate acyltransferase (GPAT). This we achieved earlier in yeast (See US Provisional Patent Application No. 60/139,788, June 21, 1999). We now report the extension of our invention to plant oils.

This is the first demonstration of an increase in carbon flux into plant triglycerides by manipulating the expression of a GPAT enzyme.

Detailed Description of the Invention

GPAT is the first enzyme of membrane and storage lipid (triacylglycerol; TAG) biosynthesis in living organisms. It determines the type of fatty acid incorporated in the sn-1 position of membrane and storage lipids. Overall, its specificity for various acyl-CoA appears to be flexible (Frentzen, 1993; Frentzen and Wolter, 1998). Since it is the first step in triglyceride and phospholipid lipid biosynthesis, it is surmised that enhancing production of GPAT might also result in an increase of 1-acylglycerol-3-phosphate through the Kennedy pathway, resulting in higher oil content. Three types of GPAT have been reported from plants: plastidial (P), mitochondrial (M), and cytosolic (ER). They exhibit variable specificity toward acyl-CoA and acyl-ACP substrates (Frentzen, 1993). The ER form of GPAT is considered to be the most important for TAG manipulation but it has not been possible to isolate a gene for this enzyme. However, cDNAs for P-GPATs has been isolated and characterized from a number of species. Also, heterologous GPAT genes from bacteria and mammals are available (see Wilkison and Bell, 1997; Dircks and Sul, 1997).

A plastidial GPAT gene has been used by others to change the fatty acid composition of membrane lipids and to improve chilling tolerance (Murata, 1992 and Nishizawa, 1996). The bacterial GPAT gene (*plsB*) has also been used to change the fatty acid composition of membrane lipids and to decrease chilling tolerance (Wolter, 1992). None of the studies thus far have used GPAT genes to increase oil content in a living organism.

Because of the unavailability of an ER GPAT gene from plants, we set out to use a plant plastidial GPAT and a bacterial GPAT gene to establish the proof of our concept. Here we demonstrate the use of a plant plastidial or bacterial GPAT gene to increase oil content in plant seeds. Two genes were modified to create four versions to achieve higher oil content.

A cDNA (-gene) for a P-GPAT has been isolated in this lab from safflower (*Carthamus tinctorius*) (Bhella and MacKenzie, 1994). A P-GPAT can use both acyl-ACP and acyl-CoA substrates, although the latter with lesser efficiency (Frentzen, 1993). It was surmised that in the absence of any acyl-ACP in the cytosol, a P-GPAT will have to use acyl-CoA, although it has to compete with the ER-GPAT for substrates, acyl-CoAs and glycerol-3-phosphate (G-3-P). Its competitive ability could be enhanced by overexpressing it by using a strong constitutive or a tissue specific promoter. Normally, a P-GPAT is targeted to plastids by a transit peptide (TP). The TP was deleted to confine the GPAT activity in the cytosol (P-GPAT-TP). The modified gene was labelled as *ctpgpat-tp*. It is anticipated that an ER environment may also be required for a P-GPAT-TP to effectively participate in TAG biosynthesis. The ER retention signals (ERRS; Jackson et al., 1990) were added to P-GPAT-TP to meet this requirement. The modified gene was labelled as *ctpgpat-tp+errs*.

The *Escherichia coli*'s *plsB* gene has been previously characterized (see Wilkison and Bell, 1997). The *plsB* can use both acyl-ACP and acyl-CoA equally effectively. The *plsB* gene was used as such and also with an added ERRS sequence. The gene constructs were labelled as *plsB* and *plsB+errs*.

EXPERIMENTAL

Preparation of chimeric genes and expression vectors

An open reading frame (*orf*; ~1.1 kb) without *tp* was amplified by PCR from the *ctpgpat* cDNA. Another chimeric gene containing an *err*s at its 3' end was also PCR amplified. The *orf* of the *E. coli plsB* gene (~2.5 kb) was PCR amplified in unmodified form or with an *err*s at its 3' end. The blunt-end PCR fragments generated using *Pfu* DNA polymerase were cloned into pSK II cloning vector. The genes were labeled as *ctpgpat-tp*, *ctpgpat-tp+err*s, *plsB* and *plsB+err*s. The intact *ctpgpat* cDNA was also used (Bhella and MacKenzie, 1994). Their sequences are shown in, respectively, Figures 1 to 5. It should be noted that the sequences representing the *ctpgpat* are different from that published earlier (Bhella and MacKenzie, 1994). There were errors in the published sequence.

For plant expression study, chimeric genes were recovered from the cloning vector and cloned into the plant transformation vector, pHS737 (landem 35s CaMV promoter) for constitutive expression of GPATs. The recombinants were transformed into *Agrobacterium tumefaciens* GV 3101 for *Arabidopsis* transformation.

Transformation of *Arabidopsis thaliana*

Arabidopsis thaliana plants were transformed by the floral dip method. Seeds (T_1) from these plants were collected and selected on a growth medium containing kanamycin. Transgenic plants were grown to maturity and seeds (T_2) from 10 individual plants were collected and used for lipid analysis. Wild type and plants transformed with vector alone were grown as controls along with the transformed plants.

Lipid analysis

The fatty acid composition of the seeds was determined by GC analysis following extraction of the oil and conversion of the triglycerides to fatty acid methyl esters. A known amount of C15 triglyceride was added to the seed sample as a marker before oil extraction. Total seed lipid content was estimated on the basis of the recovery of C15 fatty acid methyl ester.

RESULTS AND DISCUSSION:

Most of the transgenic plants appeared normal in morphology. Those that were not were discarded. The oil compositions and seed sizes of selected samples representing each construct are shown in Table 1 and Figure 5. Seeds of plants transformed using only the pHS737 vector were indistinguishable in oil content from wild type control plants grown under the same conditions. All other gene constructs produced higher seed oil content. The unmodified *spgpat*, which is expressed in the plastid, produced oil increases ranging from 10 to 21%, suggesting that lysophosphatidic acid is released from the plastids. On average the greatest increase in oil was observed in seeds of transformants carrying the *spgpat-tp* gene (average +22%). Addition of an ER targeting sequence had no apparent effect on seed oil content. The *plsB* gene increased seed oil content by an average of 15%. The addition of an ER targeting sequence resulted in an average seed oil increase of 18% but the difference cannot be regarded as significant. It is somewhat unexpected that the *spgpat* protein, which ordinarily acts on an acyl-ACP substrate and has a much reduced activity with acyl-CoA substrates, effected a greater seed oil increase (on average) than the *plsB* protein. The *plsB* protein reacts equally well with acyl-ACP and acyl-CoA substrates.

Seeds of plants transformed with the vector only did not differ significantly in average weight from wild type plants (Table 1). Seeds of individual plants from each construct were significantly larger than wild type and the pHS737 control; e.g. 315-2, 301-2, 302-6, 303-3 and 304-15. However, increased seed oil content was not always positively correlated with increased seed weight; e.g. 303-7 and 304-1. Samples representing *spgpat* genes only were more highly correlated (coefficient 0.53) than those representing *plsB* genes (coefficient 0.18).

The phenotypes representing increased seed oil content and size would result in increased yields in oilseed crops. Those representing an increase in seed oil content without an increase in seed size would provide more oil per tonne of seed and would present advantages to oilseed processors. Appropriate phenotypes can, therefore, be chosen for specific applications.

We have demonstrated that expression of both a bacterial and a plant GPAT increases plant seed oil content and seed size.

Some compositional changes were also observed in the seed oils of the transformants. Selected examples are illustrated in Table 2. Seed oils from plants transformed with the vector alone (pHS737) were not significantly different from the untransformed wild type *A. thaliana*. No significant changes were observed in the proportions of any of the saturated fatty acids or of 18:2 and 20:1. However, the proportions of 18:1 Z9 (oleic acid) decreased and of 18:3 Z9, Z12, Z15 (α -linolenic acid) increased in some of the individual transformants of all constructs. Examples are 315-6, 315-8, 301-2, 302-5, 303-3 and 304-4. These compositional changes do not always coincide with an increase in total seed oil content; for example, 315-6. Furthermore, increases in seed oil content are not always accompanied by a change in composition; for example 315-4 and 303-7.

We hypothesize that overexpression of GPAT increases the proportion of lyso-phosphatidic acid which contains oleic acid at the *sn*-1 position and, therefore, in turn, the proportion of phosphatidic acid having the same *sn*-1 fatty acid composition. The latter constitutes the substrate for subsequent desaturation by the Δ -12 fatty acid desaturase to produce linoleic acid which is subsequently further desaturated by the Δ -15 fatty acid desaturase. The result is increased proportions of α -linolenic acid, the terminal product of the desaturation pathway.

It is surprising that the change in the proportions of oleic and α -linolenic acids is observed for both the safflower and *E. coli* GPATs. In *in vitro* assays, the safflower plastidial GPAT prefers unsaturated fatty acids as compared to the *E. coli* GPAT which prefers saturated fatty acids. Nevertheless, both genes, in modified and unmodified forms, present an approach to decreasing the proportion of oleic acid and increasing the proportion of linolenic acid in contexts where such a result is desirable. In the latter context, the process is independent of any intellectual property rights vested in any of the fatty acid desaturases.

Since GPAT is a ubiquitous enzyme for phospholipid and triglyceride biosynthesis and the biochemical pathway is identical in all living organisms, it is reasonable to expect that the overexpression of these genes will also enhance oil synthesis in other microorganisms and other plant species. It is also reasonable to expect that expression of other GPATs will have similar effects on seed oil content.

Summary

An increase in triglyceride synthesis and average seed size following expression in *Arabidopsis thaliana* of an *E. coli* GPAT and a modified safflower GPAT has been demonstrated.

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Table 1. Oil content of wild type and transformant *Arabidopsis thaliana* seeds

Sample	Transformant	Oil Content		WT/100 seeds (mg)
		(Wt %)	% Increase	
Wild type	Average (n=7)	26.9 ± 0.8		1.42 ± 0.29
<i>pHS737</i>	Average (n=7)	26.8 ± 1.09		1.41 ± 0.26
<i>ctpgpat</i>	315-2	32.4	20.9	2.47
	315-3	29.4	9.7	1.92
	315-4	30.1	12.3	1.90
	315-7	29.6	10.4	1.85
<i>ctpgpat - tp</i>	301-2	34.0	26.9	2.03
	301-3	34.6	29.1	1.90
	301-5	29.3	9.3	1.79
<i>ctpgpat-tp+errs</i>	302-2	29.4	9.7	1.44
	302-6	32.7	22.0	2.45
<i>plsB</i>	303-2	29.8	11.2	1.28
	303-3	30.8	14.9	1.98
	303-4	29.1	8.6	1.89
	303-7	33.2	23.9	1.53
<i>plsB + errs</i>	304-1	30.9	15.2	1.51
	304-2	32.5	21.3	1.60
	304-3	30.3	13.1	1.38
	304-15	32.5	21.3	2.20

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Table 2. Fatty Acid Composition of Wild Type and Transformed *Arabidopsis thaliana*

Sample/construct	Wt % Oil	18:1	18:3
Wild type	26.9 ± 0.8 (n=7)	14.4 ± 0.7	15.7 ± 0.8
pHS737	26.8 ± 1.1 (n=6)	12.4 ± 1.3	15.9 ± 0.4
spgpat			
315-2	32.4	13.6	18.4
315-4	30.1	14.7	16.5
315-6	25.2	9.1	18.4
315-8	27.1	9.2	19.9
315-10	28.3	12.4	17.5
spgpat-tp			
301-1	27.5	12.0	17.1
301-2	34.0	10.9	18.8
301-3	34.6	12.2	18.2
spgpat-tp+errs			
302-4	28.1	12.5	18.5
302-5	25.6	12.2	17.5
302-6	32.7	12.9	17.5
302-7	27.8	15.8	16.5
plsb			
303-1	28.4	12.0	17.8
303-3	30.8	11.4	17.6
303-4	29.1	14.7	17.1
303-7	33.2	14.8	15.9
plsb + errs			
304-2	32.5	12.4	17.0
304-3	30.3	12.1	16.8
304-4	23.5	10.7	18.0
304-9	27.4	13.9	17.4
304-15	32.5	14.1	16.9

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Figure 1. Nucleotide sequence of the gene *ccpgpat1* (*Carthamus tinctorius* plastidial) used for transforming yeast and *Arabidopsis thaliana*. The start and stop codons are underlined. Nucleotides that are different from the published sequence are highlighted.

[illegible]

The translated protein sequence of the gene *ctgpgpat* used for transforming yeast and *Arabidopsis thaliana*. Amino acids that are different from the published sequence are highlighted.

SLSLTHNTQNTYHCYFLSLLLHSPFLAMS IFFSPSPSTLFFSTTNNANRPVSPSSSPSSAFTTPLLSSSLRLPILRGFFCLA
FSPANAAHGTAETVHGKNKWPSPSSSSAATQSPAGSDGHSHTFIDARESEQULLSGIUREAGTLPKHIAQMHAE
QNYKNAVLGSAAPHAEDIVLSNMVAFDRMLDVKEFFEFSPYHEALLERFNMYMFGNYIREFLVNFRFSPVQNVSV
FVGESEQLKQGDUNVILSNHQTADPAIVIAIMLETTNHHI SENTIYVAGDVRITDPLCKSSMGRNLLCVYSKXHMND
VPELAEMKKRSNTSLKGLMALLRGSKIIWTAPSGGHRDPTITNQWPAFVDTSLTNNMRLVODHAGLVYHITPLA
ILCHDIMPPLQVEKEIGENHISFHTGTGISVAPEINFQEVTAACSGSFEEAKAAYSQALYDSVCEQYKVLHSAVHGGK
GLEASTPSVLISOPLOFLD

Figure 2. Nucleotide sequence of the gene *ctpgpat-tp* (*Carthamus tinctorius* plastidial GPAT from which the sequence for a putative transit peptide was removed) used for transforming yeast and *Arabidopsis thaliana*. The start and stop codons are underlined. ATG was introduced as a start codon in place of GAC. Nucleotides that are different from the published sequence are highlighted.

GGATCCATGCACGGTCACTCTCGTACATTCATCGATGCTCGTTCCGAACAAGATCTTCTTTCTGGAATTCAAAGAGAG
TTGGAAGCTGGAAACACTGCCAAAACATATTGCTCAAGCAATGGAGGAGCTATATCAGAAGTACAAAAATGCAGTTCTC
CAAAGTGGCGCTCCTCATGCAGAGATATTGTGTGCAAAACATGCGTGTAGCGTTTGATCGTATGTTCTTGGATGTG
AAGGAGCCGTTTGAATTTTCAACCATATCATGAAGCTATTTTGGAACTTTTAACTACTATATGTTTGGTCAAATTTAT
ATTCCGCCCTTTGCTCAATTTCAAGGAATCATACGTTGGCAATGCTCTCCGTTTTCGGTGTATGGAAGAGCAGCTTAAG
CAGGGTGACAAGGTGGTTTTGATCTCAAACCATCAAACAGAGCAGATCCAGCTGTTATTGCCTTGATGCTTGAACA
ACAAACCCCATATTTCTGAGAACATAATCTACGTGGCAGGGGATAGACTAATAACAGATCCTCTTTGCAAGCCTTTC
AGCATGGGAAGGAATCTGTTGTGCGTGTATTCAAAAAGCATATGAATGATGTTCTCTGAGCTTGCTGAGATGAAAAA
AGATCAAATACAAGAAGTTTAAAGGCTTGGCTTTGCTTTTGGAGGGCGGATCTAAATATATGGAATTGCGCCAAGT
GGTGGCAGGGACAGGCCAGATCCTATCACAAATCAGTGGTTTCCGGCACCGTTTGATGCCACTTCGCTTGACAACATG
AGAAGGCTCGTGGACCATGCTGGTTTGGTGGGTCAATATATCCTTTAGCCATATTTGTGCATGACATCATGCCCCCT
CCTCTTCAGGTTGAGAAAGAAATGGAGAGAAGAGGATCTCTTTTATGGCACCGGAATATCAGTGGCACCGGAR
ATTAATTTCCAAGAAGTTACTGCTCTTCTGGGTCCCCCGAGGAGGCGAAGGCAGCTTATTCACAGGCATCTATGAT
TCCGTGTGTGAACAATACAAGGTGCTACATCTGCGGTACATGGAGGAAAAGGTTAGATSCATCAACACCAAGTGTG
TCGTTGTCACAACCCCTTGCACTTTCTCGATTAGGATCC

The translated protein sequence of the gene *ctpgpat-tp* used for transforming yeast and *Arabidopsis thaliana*. The mature polypeptide will not have the first 90 amino acids of the *ctpgpat* (Bhalla and MacKenzie, 1994). Amino acid 91, Asp (D), is replaced with Met (M). Amino acids that are different from the published sequence are highlighted.

GSMDHSRTFIDARSEQDLLSGIQRELEAGTLPKKIAQAMEELYQNYKNAVILQSAAPHAEDIVLSNMRVAFDRMFLDV
KEFFEFSPYREAILPEFNYMFGQNYIRPLVNFRESYGVNVSFVGMEEQLKQGDVVLISNHTQADPAVIALMLET
TNPRISENIIVAGDRVITDPLCKPFMSGRNLLCVYSKKNNDVPELAEMKKRSNTRS1KEMALLLRGGSKIWIAPS
GGRDRPDPIITNQWFPAPFDATSLDNMRRVLVDHAGLVGHYIPLAILCHDIMPPLQVEKLIGEKISFHTGTGISVAPE
INFQVNTASCCSPPEEAKAAYSQALYDSVCBQYKVLHSAVHGGKGLEASTPSVLSQPIQFLD.D

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Figure 3. Nucleotide sequence of the gene *ctgpat-tp+err*s (*Carcharias tinctus* plastidial GPAT from which the sequence for a putative transit peptide was removed and sequences for endoplasmic reticulum retention signals were added). Start and stop codons are underlined. ATG was introduced as a start codon in place of GAC. Additionally, AAG and AAA (also underlined) in place of TTG and TTT were introduced for ER retention of the protein. Nucleotides that are different from the published sequence are highlighted.

GGATCCATGCACGGTCACTCTCGTACATTATCGATGCTCGTCCGAAACAGATCTTCTTTCTGGAATTCAAAGAGAG
TTGGAAGCTGGAACACTGCCAAAACATATTGCTCAAGCAATGGAGGAGCTATATCAGAACTACAAAAATGCAGTTCTC
CAAAGTGCGGCTCCTCATGCAGAAGATATTGTGTGTTCAACATGCGGTAGCGTTTGATCGTATGTTCTTGGATGTG
AAGGAGCCGTTTGAATTTTACCATATCATGAAGCTATTTTGAACCTTTTAACTACTATATGTTTGGTCAAAATTAT
ATTGCGCCTTTGGTCAATTTCAAGGAATCATACGTTGGCAATGTTCTCGGTTTTCGGTGTANTGGAAGAGCAGCTTAAG
CAGGGTGACAAGGTGGTTTGTGATCTCAAACCATCAAACAGAGCAGATCCAGCTGTTATTGCTTGAATGCTTGAACA
ACAAACCCCATATTTCTGAGAACATAATCTACGTGGCAGGGGATAGAGTAATAACAGATCCTCTTTGCAAGCCTTTC
AGCATGGGAAGGAATCTGTTGTGCGTGTATTCAAAAAGCATATGAATGATGTTCTCTGAGTTGCTGAGATGAAAAA
AGATCAAAATACAAGAAGTTTAAAGATTTGGCTTTGCTTTTGAAGGGCGGATCTAAATAATATGGATTGCGGCAAGT
GGTGGCAGGGACAGGCCAGATCCTATCACAAATCAGTGGTTTCCGGCACCGTTTGATGCCACTTCGCTTGACAACATG
AGAAGGCTCGTGGACCATGCTGGTTTGGTGGGTCAATATATCCTTAGCCATATTGTGCCATGACATCATGCCCTCT
CCTCTTCAGGTTGAGAAAGAAATTGGAGAGAAGAGATGATCTCTTTTCATGGCACCGGATATCAGTGGCACCGGAA
ATTAATTTCCAAGAAGTTACTGCTCTTGTGGGTCCCCGAGGAGGCGAAGGCAGCTTATTCACAGGCACTCTATGAT
TCCGTGTGTGAACATAACAAGGTGCTACATTCTGCGGTACATGGAGGAAAGGGTTAGAGCATCAACACCAAGTGTG
TCGTTGTCAACCCCAAGCAGAACTCGATTAGGATCC

Translated protein sequence of the gene *ctgpat-tp+err*s. The mature polypeptide will not have the first 90 amino acids of the *ctgpat* (Bhella and MacKenzie, 1994). Amino acid 91, Asp (D), is replaced with Met (M). Also Lys (K), underlined, replaced a Phe (F) and Leu (L) at position 3 and 5 from the C terminus. Amino acids that are different from the published sequence are highlighted.

GSMHGHSTFIDARSEQDLLSGIQRELEAGTLFKRIQAAMEELYQNYNAVLQSAAPHLEDIVLSNMRVAFDRMFLDV
KEFFEFSFYHEAILEFPNYMFGQNYIRPLVNFRESYVGNVSVFGVMEELKQGDVVLISNMQTEADPAVIAIMLET
TNPHISENIIVAGDRVITDPLKPFSSMGRNLLCVYSKKMNDVPELAEMKKRSNTRSLKQALLRGGSKIWIWAPS
GGDRDPDITNQWFPAPFDATSLDNMRRLVDHAGLVGHIYPLAILCHDIMPPLQVEKEIGEXKISFHGTGISVAPE
INFQEVITACGSPBEAKAAYSQALYDSVCEQYKVLHSAVHGKGLEASTFVSLSQPKQLD.D

Figure 4. Nucleotide sequence of the gene *plsB* (*E. coli* GPAT). Start and stop codons are underlined. A nucleotide that is different from the published sequence is highlighted. It was introduced during PCR.

AGATCTTCCCATGACTTTCTGCTATCCTTGGCCGCCATTGTCATTATTAAACAGAGGCTTACATCGTTTATGTCCGG
CTGGCCACGAATTTACTACAAATTACTGAATTTACCATTAAGCATCCTGGTAAAAAGCAAGTCTATTCCGGCAGATCC
TGCCCGGAACTGGGGCTGGATACCTCTCGTCCAATTATGTACGTTTTACCGTACAACTCGAAAGCAGATTGTGTAC
GTTGCGCGCCAGTGTCTGGCACATGACTTGCTGACCCGTTAGAGCCGCTGGAAATCGACGGCCAGCTACTGCCCGG
CTATGTGTTTATTACGGCCGGGCGCGTGTGTTACCTATTACAGCCGAAAGAGAGTCTATTAGCTGTTCCACGA
GATCTCGATTTGCACCGTAGCAACCAAACTGCGATGTGCAATGGTCCAGTGTCCGTTATGTTGGTCCGCGCC
GGGCGGTGAAAAAGGCGAAGTGAACCCCGCGCTGCGTATGCTTAACGGCGTACAGAAATTTTCGCTGTACTGTGGCT
CGGTCCGACAGTTTTGTGCGTTTCTCGCCGTCAGTTTCGCTGCGCGGTATGGCGGATGAACACGGCCAGGATAAAAC
TATCGCTCAGAACTGGCGCGCGTGGCGCGTATGCACTTTGCCCGTCAACGTCTGGCTGCCGTAGGCCACGTTTGGC
TGCTCGTCAGGATCTGTTAATAAGCTGCTCGCTCCCGCGCCATTGCCAAAGCGGTAGAGATGAAGCCGCGCAGCAA
AAAAATCTCCCATGAAAAAGCCGAGCAGAACCGGATTGCACTGATGGAAGAGATTGCGGCGAATTTCTTACGAGAT
GATTCCGCTGACTACCGTATTCTGGGCTTCACTGGAACCGACTTTACAGGGCATCAACGTCATACCGTACGCGG
TTCCTACGTTGCTGATCACCAGGGGCTGTTGCCCGCGCATATCGCCCGCGGATCAACCTGAATTTCTGGCTGCCG
GCGGATTTTCGCGCGTCTGGGGCGCTCTTTTATTCGCGTACGTTTAAAGGCAATAAACTTTATTCACCGTTTTCG
GGGATCTCGGCGAATCTTTCAGCCGTTGGTTATTCCGTCGAGTACTTCGTTGGAAGGCGTTCGTTCCCGTACGGGGG
TTTGTGGATCCGAAACTGTTACGCTGTGCGATGACATTACGGCGATGCTCCGTCGGCGCAGCGCTCCGATTACGCT
GATTCCGATCTATATCGGTTATGAGCAGCTCATGGAAGTGGGTACTTACGCCAAAGAACTGCGCGCGCGCAGCAAGA
GAAAGAGAGCTTCCCGCAGATGCTGCGCGGTTTAAAGCAAGCTGCGTAACTCTCGGTGAGGTTACGTCAACTTCGGTGA
ACCAATGCCGTTGATGACCTACCTTAAACAGCATGTACCTGACTGGCGTGAATCTATCGATCCCATCGAAGCGGTGCG
TCCGGCATGGTTAAGCCGACGGTCAATAATATGCTGCCGATCTGATGGTACGCATTACCAACGAGCGCGCGCAAA
CGCCATGAACCTGTGCTGACTGCGCTACTGGCATCAAGTCAGCGCTCACTACCCCGCAGCAGTTAACCGAGCAACT
CAACTGCTACTGGATCTGATGCGCAACGTTGCCCTACTCCACGACTCTACCGTTCCCTCAGCCAGCGCCAGCGAGT
TATCGATCAGCGCTGCAAAATGAACAAGTTTGAAGTCGAGAAAGACACAATCGGCGACPTCATCATCTGCGCGCGCA
GCRAGCGGTGCTGATGACCTACTATCGCAACACATTGCGCATATGTTGGTGTGCTTCGCTGATGGCGCAATCGT
CACCCAGCATCGCCACATCTCCCGCAGCTATTGATGAGCAGCTCAATGTGCTTTACCCAATGCTGAAAGCGGAGCT
GTTCTGCGCTGGGATCGCGACGAGTTGCCCGACGTTATTGATGCGCTGGCAATGAGATGCAACGTACGGGCTGAT
TACCTGCAAGATGATGAGTTGCATATCAACCCGCGCGCTCTCGACCGCTACAGCTGTGGCCGCGAGGCGCGCGCA
AACGCTGCAAGCTTATGCCAICACCTTCTGGTTGTTGAGTGGCCAAACCGCTCGATCAACGCGGTACGCTGAGAGA
GAGCCGACCGTCCGCAACGCTCTCCGCTGCTGCACGGCATCAACGCGCGGAGTTCTTCGACAGGCGGTGTTTCAG
TTCTCTGGTGTGACTGCGTATGAAGGATATATCAGCGATAGCGCGGATGCCGAACCGGCGAGAACGATGAAGGT
TTATCAGTTGCTGGCGAGTTGATTACATCAGACGTCGCTTGACGATTGAGAGTGCGACGCGAGGCGGAGGTTATC
AGATCT

Translated protein sequence of the gene *plsB* (*E. coli* GPAT). An amino acid that is different from the published sequence is highlighted.

MTFCYPCRAFALLTRGFTSFMSGWPRIYYKLLNPLSLVSKSIPADPAPELGLDTHRPIMXVLPYNSKADLLTLRA
QCLAHDLPLDPLEIDGTLPLRYVFIHGGPRVFTYTPKEESIKLFHDYLDLHRSNPNLDVQMVVSVMFGRAPGRE
KGEVNEPLRLNLGVQKFFAVLWLGRDSFVRFSFVSILRRMADEHGTDKTIAQKLARVARMHFARQRLAAVGPRLPARQ
DLFNKLLASRAIAKAVEDEARSKKISHEKAQQAIALMEEIAANFSYEMIRLTDRLGFTWNRLYQGIVINAERVRO
LAHDGHELUVPCERSHMDYLLLSYVLYHQGLVPPHIAAGINLNFWPAGPIFRRLGPFIRRTFKGNKLYSTVREYIL
GELFSRGYSVEYFVGGRSRTGRLLDPKGTGLSMTIQAMLGGTRPITLPIYIYGYHVMVEVGTYAKELRGATKES
LPQMLRGLSKLRNLGGYVNFGEPMELMTYLNQHVDPWRESIDPIEAVRPAHLTFTVNNLAADLMVRINNAGAAANAMN
LCCTALLASRQSLTREQLTEQLNICYLDLNRNVPYSTDSTVPSASASELIDHALQMAKFEVEKDTIGDIIILPREQAV
LMTYRNINIAHMLVLPSSMAHIVTQHRHISRDLMEHVNVLPMKAEFLRWDRDFLPDVIDALANEMQRQGLITLQ
DDELHINPAHSRTLQLLAAGARETLQRYAITFWLLSANFSINRGTLKESRTVAQRJ.SVLHGINAPEFFDKAVFSSLV
LTLRDEGYISDSGDAEPAETMKVYQLLAEILTSVRLTIESATQGG.SD

Figure 5. Nucleotide sequence of the gene *plsB+errs* (*E. coli* CPAT to which sequences for endoplasmic reticulum retention signals were added). Start and stop codons are underlined. AAG and AAG (also underlined) in place of GCC and ACG were introduced at the 3' end for ER retention of protein.

AGATCTTCCCATGACTTTCTGCTATCCTTGCCGCGCATTTGCATTATTAACCAGAGGCTTTACATCGTTTATGTCCGG
CTGGCCACGAATTTACTACAAATTACTGAATTTACCATTAAAGCATCCTGGTAAAAAGCAAGTCTATTCGGGCAGATCC
TGCCCGGGAAGTGGGGCTGGATACCTCTCGTCCAAATTATGTACGTTTACCGTACAACTCGAAAGCAGATTGTCTGAC
GTTGGCGGCCCACTGTCTGGCACATGACTTGCTGACCCGTTAGAGCCGCTGGAATCGACGGCACGCTACTGCCGCG
CTATGTGTTCAATTCACGGCGGGCGCGTGTGTTCACCTATTACACGCCGAAGAAGAGTCTATTAAAGCTGTTCCACGA
CTATCTCGATTGACACCGTAGCAACCCAAATCTGGATGTGCAGATGGTGCCAGTGTCTGGTGTATGTTGGTTCGCGCGC
GGGGCGTGAAAAAGCGAAGTGAACCCGCGCTGCGTATGCTTAACGGCGTACAGAAATTTTCGCTGTACTGTGGCT
CGGTCCGCACAGTTTGTGCGTTTCTCGCCGTGAGTTTCTGCTGCCGCTATGGCGGATGAACACGGCACGGATAAAAC
TATCGCTCAGAAATCGGCGCGCGTGGCGCGTATGCACTTTGCCCGTCAACGTCTGGCTGCCGTAGGCCCCAGTTTGGC
TGCTCGTCAGGATCTGTTAATAAGCTGCTCGCCTCCCGCGCATTTGCCAAAGCGGTAGAGATGAAGCGCGCAGCAA
AAAAATCTCCCATGAAAAAGCGCAGCAGAACGCGATTGCACTGATGGAAGAGATTGCCGCGAATTTCTCTTACGAGAT
GATTCGCTGACTGACCGTATTCTGGGCTTCACTGGAACCGACTTTACAGGGCATCAACGTCCATACCGCTGAGCG
CGTTCCGCGAGCTGGCCACGACGGTCAATGAGCTGGTATATGTGCTTGGCCACCGCAGTCAATGGACTACCTGCTGCT
TTCTTACGTGCTGTATCACCAGGGCGCTGGTGGCGCGCATATCGCGCGCGGATCAACCTGAATTTCTGGCTTCCGCG
CGCGATTTTCGCGCGTCTGGGGCGGTTCTTATTCGCGGTACGTTTAAAGGCAATAAACTTATTCCACCGTTTTCG
GGAGTATCTCGGCGAATCTGTTACGCGGTGGTTATTCGCTCGAGTACTTCTGGAAGCGGTCTCTTCCCGTACGGGGCG
TTTGTGATCCGAAACTGGTACGCTGTGATGACCATTCAGGCGATGCTGCTGGCGGCACGCGTCCGATTACGCT
GATTCGGATCTATATCGGTTATGAGCACGTCATGGAAGTGGGTACTTACGCCAAAGAACTGCGCGCGCGACGAAAGA
GAAAGAGAGCCTGCCCGCAGATGCTGCCGCGTTTAAAGCAAGCTGCGTATCTCGGTGAGGTTACGTCAACTTCGGTGA
ACCAATGCCGTTGATGACCTACCTTAACAGCATGTACCTGACTGGCGTGAATCTATCGATCCCATCGAAGCGGTGCG
TCCGGCATGGTTAAGCGCGACGGTCAATAATATTGCTGCCGATCTGATGGTACGCATTAAACAGCAGGCGCGGCAAA
CGCATGACCTGTGCTGACTGCGCTACTGGCATCAGCTCAGCGCTCACTACCCGCGAGCAGTTAACCGAGCAACT
CAACTGCTACCTGGATCTGATGCGCAACGTGCCCTACTCCACGGACTCTACCGTTCTTTCAGCCAGCGCCAGCGAGCT
TATCGATCAGCGCTGCAAAATGAACAAGTTTGAAGTCGAGAAGACACAATCGGCGACATCATCATTTCTGCCGCGCGA
GCAAGCGGTGCTGATGACCTACTATCGCAACAACATTGCGCATATGTTGGTGTCTGCTTCGCTGATGGCGGCAATCGT
CACCAGCATCGCCACATCTCCCGCGAGTATTGATGGAGCAGCTCAATGTGCTTTACCCAACTGCTGAAGCGGAGCT
GTTCTGCGCTGGGATCGCGACGAGTTGCCGCGGCTTATGATCGCTGGCAATGACATGCAACGTGAGGGGCTGAT
TACCCTGCAAGATGATGAGTTGCATATCAACCCGCGGCTTCTCGCACGCTACAGCTGCTGGCGCGCAGCGCGCGGA
AAGCGTGCACAGCTTATGCCATCACCTTCTGGTGTGATGAGTGCCAAACCGTTCGATCAACCGCGGTACGCTGGAGAAAGA
GAGCCGACCGCTCGCGCAACGTCTCTCCGTGCTGCACGGCATCAACGCGCGGAGTTCTTCGACAGGCGGTGTTTCAG
TTCTCTGGTGTGACACTGCGTGATGAAGGGTATATCAGCGATAGCGCGATGCCGAACCGCGAGAACGATGAAGGT
TTATCAGTTGCTGGCGGAGTTGATTACATCAGACGTGCGTTTGACGATTGAGAGTGCAGAGCAGAGGAAGGTAATC
AGATCT

Translated protein sequence of the gene *plsB+errs*. Amino acid Lys (K), underlined, replaced a Gly (G) and Thr (T) at position 3 and 5 from the C terminus.

MTFCYPCRAFALLTRGFTSEMSGWFRYYKLLNLPLSLVKSISIPADPAPELGLDTSRPIYVLPYNSKADLLTLRA
QCLAHDLPLEPLEIDGTLPLRYVFIHGGRFVFTYTPKEESIKLFHDYLDLHRSNPNLDVQMPVSVMFGRAPGRE
KGEVNPPLRLNGLVQKFFAVLWLGKDSFVRFSVSLRRMADEHGTDKTIAQKLARVARMHFAQRILAAGVPRLPARQ
DLFNKLASRAIAKAVEDEARSKKISHEKAQONAIAMEEIAANFSYEMIRLTDRIILGFTWNRLYQGINVHABRVQ
LAHDGHELVYVPCHRSHMDYLLSYVLYHQGLVPPHIAAGINLFWPAGPIFRRLGAFPIRRTFKGNKLYSTVFREYL
GELFSNGYSVEYFVEGGRSRTGRLLDPKGTLSMTIQAMLRGGTRPITLPIYIYGEHVMEVGTYAKELRGATKEKES
LPQMLRGLSKLRNLGQGYVNFGEPMPLMTYLNQVFDWRESIDPIEAVRPAWLTPTVNNIAADLMVRINNAGANAMN
LCCTALLASRQRLTREQLTEQLNLYLDMRNVPYSTDSTVPSASASELIDHALQHNKFEVEKDTIGDIILPREQAV
LMTYRNINARMLVPLSMAAIVTQRHISRDLMERHNVLYPLKAEFLRWDRDELDPVIDALANEMRQGLITLQ
DDELHINPABSRITQLLAAGARETLQRYAITFWLLSANPSINRGTEKESRTVAQQLSVLHGINAPEFFDKAVFSSLV
LTLRDEGYSISDGAEPATMKVYQLLAELITSDVRLTIESAKQKEG

009120-50629109

Figure 6. Seed Oil Content of Wild Type and Transformed *A. thaliana*